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Chapter 7

Genomics of Tropical Fruit Tree Crops

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Abstract The genetic improvement of tropical fruit trees is limited when compared to progress achieved in temperate fruit trees and annual crops. Tropical fruit tree breeding programs require significant resources to develop new cultivars that are adapted to modern shipping and storage requirements. The use of molecular markers in tropical fruit tree breeding is greatly assisting in solving a number of difficult challenges for breeders such as the development of complex family structures for recombination mapping and for recurrent selection. A review of the literature on molecular markers development and new techniques for increasing single-nucleotide polymorphic markers is discussed. The development of marker-assisted breeding for these tropical tree crops is also discussed.

Keywords SNP • Microsatellites • SSR • Molecular markers • Mango • Avocado • Lychee • Longan • Linkage mapping

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Introduction

Genetic improvement of tropical fruit trees¹ has lagged far behind the progress achieved in temperate fruit trees. The reasons for this are many and include the lack of resources, political stability, and infrastructure in many of the lesser-developed countries where these species are grown. Nevertheless, the value of these tree fruit crops in providing locally available quality nutrition is significant. The commercial potential of these crops is often limited because of logistics involving transportation and storage. The commercial value of these crops could be greatly enhanced if cultivars that were adapted to commercial production, storage, and shipping were developed.

Tropical fruit trees are a major source of carbohydrates and vitamins for much of the developing world population. The world production of the major tropical fruits from trees is expected to reach 65 million tons for the year 2010, with 98% of the global production by developing countries. The production in the year 2010 for the top three tropical fruit trees, mango, papaya, and avocado, is expected to be 30.7, 12.4, and 3.1 million tons, respectively. For the minor tropical fruit trees, such as lychee, durian, rambutan, and guava, the production is smaller, and trading occurs at regional markets; therefore, there are no global statistics for these species (FAO 2003).

Compounds with a vast number of applications are present in tropical fruit trees. To mention some examples, *Garcinia mangostana* Linn. can accumulate up to 56% of oil in seeds (Hawkins and Kridl 1998). Other compounds from *Garcinia* spp. include antioxidants (Obolskiy et al. 2009) used for medicinal purposes, as well as antifungal compounds (Geetha et al. 1997) both extracted from the fruits' pericarp. The fruits of *Crataeva* spp. are high in carotenoids (Englberger et al. 2009), and the fruits of *Mangifera indica* Linn., *Carica papaya* Linn., and *Psidium guajava* Linn. are a source of carotenoids and vitamin C (Oliveira et al. 2010). Antimicrobial compounds are produced by *Diospyros blancoi* A. DC. (Ragasa et al. 2009), *Nephelium longana* Cambess. (Ripa et al. 2010), *Irvingia gabonensis* (Aubrey-Lecomte ex O. Rorke) Baill. (Kuate et al. 2007), and *Inga fendleriana* Benth. (Pistelli et al. 2009). Anticancer activity has been described for several tropical fruit tree species, for example, *Irvingia malayana* Oliv. ex A. W. Benn. has antiangiogenic properties (Ng et al. 2010), compounds from *Annona* spp. arrest cancer cells at G1 (Yuan et al. 2003), and *Morinda citrifolia* Linn. is active against cancer cells (Liu et al. 2001). Other useful applications described for tropical fruit trees are the cardioprotective activity of *Garcinia mangostana* compounds (Devi-Sampath and Vijayaraghavan 2007), the effective gastroenteritis control by *Spondias purpurea* Linn. (Caceres et al. 1993), the boost for the immune system by *Morinda citrifolia* (Palu et al. 2008), the anti-snake venom properties of *Tamarindus indica* Linn. (Ushanandini et al. 2006), and the antifatigue activity of *Dimocarpus longan* Lour. polysaccharides

¹The tropical fruit tree species discussed in this chapter will generally exclude tree species whose commercial value is derived from non-fruit parts, are predominantly temperate species, are monocots, or are discussed in another chapter in this edition.

(Zheng et al. 2010). The importance and diversity of this group of species warrant a greatly expanded effort in genetics and genomics for compound discovery and improvement of human health.

The use of genomics in tropical fruit crops has largely been confined to the development of isozyme and dominant PCR-based markers and their use for germplasm diversity analysis and clonal fingerprinting. Information from dominant markers is of limited use in genomics applications; nonetheless, it has aided in the creation of a few genetic recombination maps and in our understanding of genetic diversity in germplasm collections. More recently, codominant microsatellite markers, also known as simple sequence repeat (SSR) markers, have been developed for a number of tropical fruit crops and have been used for parentage analysis, clonal fingerprinting, genetic diversity analysis, and development of genetic linkage maps. With the increased sequencing capacity of second- and third-generation pyrosequencing, many transcriptomes (gene space) of these species will be sequenced over the next 510 years. The generation of this new information should lead to increased interest in tropical tree fruit crops and to new opportunities to increase the rate of genetic gain in breeding programs.

Breeding tropical fruit trees is complicated by their reproductive biology. For instance, avocado (*Persea americana* Mill.) has an unusual flower behavior with two complementary types of flowering patterns, called “A” and “B,” that promotes outcrossing (Bergh 1969; Davenport 1986). Avocado does not contain any self-incompatibility system, and self-pollination is frequently observed. A mature tree can produce upwards of a million flowers with only 1% of these setting fruit. A single ovary develops into the seed from a single pollination, so the generation of large numbers of seedlings by hand pollination is not practical. In mango (*Mangifera indica*), a similar situation exists; flowering is strongly influenced by weather, and some genotypes flower very irregularly. The flowers of mango are small, and both perfect and staminate (male) flowers occur in the same inflorescence. Hand pollination is possible but difficult to perform on a large scale to generate large numbers of progeny (Sharma et al. 1971 in Singh et al. 1980; Pinto et al. 2004). In addition, polyembryony in mango complicates breeding schemes. In polyembryonic cultivars, seedlings arise from nucellar tissue or from a zygote, but distinguishing between the two can be complicated (Schnell et al. 1994). For many of the tropical fruit tree species in the Sapindaceae, flower polymorphisms (i.e., monoecy, dioecy, androdioecy, gynodioecy, etc.) add complexity to the breeding process. In lychee (*Litchi chinensis* Sonn.), there are three types of flowers appearing in irregular sequence or simultaneously on the same inflorescence. There are male flowers, hermaphrodite flowers that act as females, and hermaphrodite flowers that act as males (Morton 1987). Again, the flowers are small and not amenable to hand pollination (Stern and Gazit 2003). In longan (*Dimocarpus longan*), there are three flower types: staminate (functionally male), pistillate (functionally female), and hermaphroditic (bisexual). Flowering in each panicle occurs in progressive openings of staminate (male) flowers first, then pistillate flowers followed by hermaphroditic flowers functioning as females and then hermaphroditic flowers functioning as males. Pollination is mainly by small insects but also by wind (Blanche et al. 2006).

Lack of genetic diversity in mangosteen (*Garcinia mangostana*) is a consequence of its mode of reproduction as an obligate apomict. Only anecdotal reports of male trees have been made for this dioecious species (Sand et al. 2005). In addition, an extremely long juvenile stage, upwards of 10 years, has been described for this particular fruit tree species (Poerwanto 2002).

In most of these tropical fruit tree species, controlled pollinations are very difficult to achieve, so breeders and horticulturalists have relied on the use of open-pollinated progeny for selection. The use of maternal half-sib families is very inefficient, and for most tropical fruit trees, no recurrent parentage selection has occurred. The ability to generate large numbers of codominant markers quickly using next-generation sequencing has revolutionized what can be accomplished in tropical fruit tree breeding. Identifying full-sib families is relatively simple using SSR markers. By planting trees in a polycross design (Iyer and Schnell 2009) or harvesting from commercial orchards where only a few cultivars are being grown, a breeder can now have large full-sib families to perform selection. As reported later in this chapter, the use of these types of molecular markers has greatly accelerated mango and avocado breeding efforts. Furthermore, recent efforts have focused on the development of new SSR markers for a number of tropical fruit tree species including lychee, longan, rambutan, Spanish lime, sapodilla, mamey, *Annona* spp., mangosteen, *Artocarpus* spp., and star fruit which are available to breeders and geneticists working on these crops.

Applications of Molecular Genetics and Molecular Marker Development

The main genetic tools used for the identification and breeding of cultivars of domesticated species are morphological and molecular markers (Tanksley 1993). Morphological markers in tropical fruit trees take years to be usable, for example, until the plants overcome their juvenility stage besides being subject to epigenetics and environmental factors. Molecular markers, instead, can be used from any tissue at any time during the plant growth, expediting the process of variety identification and breeding, and overcoming the limitations of traditional methods (Azofeifa-Delgado 2006). Furthermore, molecular marker analysis of tropical fruit trees help understand the past domestication of taxa, determine effective population sizes, and assess the value of a decentralized approach for future domestication (Jamnadass et al. 2009). In tropical fruit tree species for which expressed sequence tags (ESTs) are available, these can be used to develop markers such as SSRs and single nucleotide polymorphisms (SNPs) in relation to flavor, color, fragrance, vitamins, fruit softening, and other traits of interest, for example, *Actinidia* spp. (Crowhurst et al. 2008). SNPs are becoming more popular than SSRs as genetic markers in linkage analysis because they are more abundant and suitable for automatic allele calling (Novelli et al. 2004; Selmer et al. 2009). SNPs have been developed only for a few tropical fruit tree species, one in relation to resistance to papaya ringspot virus, PRSV, in *Carica papaya* Linn. (Dillon et al. 2006), another for phylogenetic studies

in *Citrus* spp. (Novelli et al. 2004) and for genotyping and linkage mapping in *Theobroma cacao* Linn. (Livingstone et al. 2010). However, at present, the cost of developing and testing SNPs is still higher than for developing SSRs. Nonetheless, SNP discovery is actively being carried out in crops such as avocado and mango.

SSRs have been the most widely employed class of molecular markers used in genetic studies with applications in many fields of genetics including genetic resources conservation, population genetics, molecular breeding, and paternity testing (Ellegren 2004). This range of applications is due to the fact that SSR markers are codominant, multiallelic, and highly reproducible; have high resolution; are amenable to high throughput; and are based on polymerase chain reaction (PCR) (Oliveira et al. 2006). As a convention, SSRs are regions in the genome where a group of bases (1–8 bp long) are repeated in tandem (Richard et al. 2008). These regions can be isolated either by data mining of existing sequences or by generating SSR-enriched libraries (Kijas et al. 1994; Zane et al. 2002). With the exception of *Carica papaya* and *Theobroma cacao* for which the genomes have been sequenced (Ming et al. 2008; Argout et al. 2010), most tropical fruit tree species do not have enough DNA sequence information to use data mining for identifying potential markers. Alternatively, expressed sequence tag (EST) information (i.e., cDNA) may be used to develop markers. We have summarized the number of entries for nucleotides, ESTs, and SSRs listed in the National Center for Biotechnology Information (NCBI) database, GenBank, for some of the most important tropical fruit trees in Table 7.1.

The use of molecular markers in tropical fruit trees can sometimes be hindered by socioeconomic reasons. One limitation is the significant cost associated with the development of markers for each crop. Tropical fruit tree species are distributed in a large number of taxonomic groups (Muchugi et al. 2008), and though transferring markers from other species could be used to reduce costs (Viruel and Hormaza 2004), such transferability is not feasible among distant taxa (Ellis and Burke 2007). In general, funds to study each tropical fruit tree species are scarce resulting in insufficient information on molecular markers. Additionally, the practical implementation of the existing results on molecular markers is often limited by the absence of guidance on how to best apply them (Muchugi et al. 2008). For example, from the top 25 indigenous tropical fruit tree species identified as priority by the International Centre for Research in Agroforestry (ICRAF), only for eight has some work been done using molecular markers (Jamnadass et al. 2009). For most of these species, the amount of genetic information available is negligible or null. A list of the main tropical fruit tree species and the molecular markers that have been developed for each of them is shown in Table 7.2.

SSR Isolation Using Pyrosequencing

The isolation of SSRs from species for which little to no genetic information is available, such as most tropical fruit trees, can be difficult. At the USDA-ARS Mid South Area Genomics Laboratory (MSAGL), an effective pipeline has been created to isolate SSR markers from these species. The process first involved a slight

Table 7.1 Summary of number of entries in NCBI for nucleotides, ESTs, and SSRs for some of the most important tropical fruit trees

Species	Nucleotide	EST	SSR	Plant type
<i>Actinidia deliciosa</i> + <i>A. arguta</i> (kiwi)	230+78	57,751+7,257	30	Vine
<i>Carica papaya</i>	51,217	77,393	45+	
<i>Musa</i> spp. and hybrids	4,210	31,268	550	Monocot
<i>Citrus</i> spp.	2,592	549,188	106	Temperate
<i>Persea americana</i>	493	16,558	0	Tropical tree
<i>Mangifera indica</i>	401	68	462	Tropical tree
<i>Cocos nucifera</i>	382	6	0	monocot
<i>Diospyros kaki</i>	296	9,474	69	Tropical tree
<i>Citrus grandis</i> (<i>C. maxima</i>)	202	0	17	Tropical tree
<i>Spondias purpurea</i>	141	0	0	Tropical tree
<i>Dimocarpus longan</i>	137	66	0	Tropical tree
<i>Annona cherimolia</i>	120	0	97	Tropical tree
<i>Inga edulis</i>	95	0	5	Tropical tree
<i>Litchi chinensis</i>	88	0	27	Tropical tree
<i>Psidium</i> sp. (guava)	75	0	24	Tropical tree
<i>Bactris gasipaes</i>	71	0	46	Monocot
<i>Durio</i> spp.	66	0	7	Tropical tree
<i>Garcinia mangostana</i>	64	149	0	Tropical tree
<i>Morinda citrifolia</i>	62	0	0	Tropical tree
<i>Adansonia</i> spp.	55	0	18	Tropical tree
<i>Passiflora edulis</i> (passion fruit)	51	0	10	Vine
<i>Manilkara zapota</i>	34	0	0	Tropical tree
<i>Averrhoa carambola</i>	31	0	0	Tropical tree
<i>Annona muricata</i>	30	0	0	Tropical tree
<i>Anacardium occidentale</i>	30	0	21	Tropical tree
<i>Tamarindus indica</i>	25	0	0	Tropical tree
<i>Irvingia</i> spp.	22	0	0	Tropical tree
<i>Nephelium lappaceum</i>	14	0	0	Tropical tree
<i>Pometia pinnata</i>	14	0	0	Tropical tree
<i>Artocarpus heterophyllus</i>	10	1	0	Tropical tree
<i>Annona squamosa</i>	8	0	4	Tropical tree
<i>Artocarpus altilis</i>	7	1	0	Tropical tree
<i>Nephelium ramboutan-ake</i>	7	0	7	Tropical tree
<i>Annona reticulata</i>	6	0	4	Tropical tree
<i>Melicoccus bijugatus</i>	6	0	0	Tropical tree
<i>Syzygium samarangense</i>	5	0	0	Tropical tree
<i>Premna serratifolia</i>	3	0	0	Tropical tree
<i>Sandoricum koetjape</i>	2	0	0	Tropical tree
<i>Garcinia portoricensis</i>	1	0	0	Tropical tree
<i>Pouteria sapota</i>	1	0	0	Tropical tree
<i>Spondias dulcis</i>	1	0	0	Tropical tree
<i>Dracontomelon vitiense</i>	0	0	0	Tropical tree
<i>Garcinia binucao</i>	0	0	0	Tropical tree
<i>Garcinia cochinchinensis</i>	0	0	0	Tropical tree
<i>Crataeva speciosa</i>	0	0	0	Tropical tree

Species were sorted first by decreasing number of nucleotide entries and then by decreasing number of ESTs. The orange highlight corresponds to species considered minor fruit crops, underutilized or rare, depending on the region, by the International Tropical Fruits Network (www.itfnet.org). The green highlight corresponds to species economically important that are either called “trees” though they are monocots, that are dicots but not exactly trees, or that are temperate

Table 7.2 Summary table of publications on molecular markers per tropical fruit tree species

Species	Work done on molecular markers
<i>Carica papaya</i> L.	CAPS marker linked to PRSV-P resistance (45); 50 accessions analyzed using nine isozyme systems (Ocampo et al. 2006); high-density genetic map with 712 SSRs and 277 AFLPs (Blas et al. 2009); draft of complete genome (97)
<i>Persea americana</i> Mill.	25 SSRs used on 37 cultivars and wild relatives (11); SNP development based on resequencing of 4 nuclear loci of 21 wild accessions of <i>P. americana</i> (Chen et al. 2008); screening of 56 SSR markers on <i>P. americana</i> (162)
<i>Mangifera indica</i> L.	15 SSR markers tested on 59 <i>M. indica</i> cultivars (137); 28 SSRs tested on 15 <i>M. indica</i> cultivars (46); 16 SSRs tested on 28 mango cultivars (157); 19 SSRs used on 307 <i>M. indica</i> accessions (Duval et al. 2006); six populations of <i>M. indica</i> analyzed by RAPD (Diaz-Matallana et al. 2009); 11 SSRs used in the phylogeographical analysis of <i>M. indica</i> (Hirano et al. 2010)
<i>Diospyros kaki</i> L. f.	Six SSRs used to screen 12 genotypes (Guo and Luo et al. 2008)
<i>Citrus maxima</i> L. Osbeck	370 accessions, 76 <i>C. maxima</i> , tested with 24 SSR markers (Barkley et al. 2006); genetic map of <i>Citrus</i> based on 256 RAPD markers used on 94 hybrids (de-Oliveira et al. 2005); putative SNPs annotated for <i>Citrus</i> (Martinez-Godoy et al. 2008)
<i>Spondias purpurea</i> , <i>S. dulcis</i>	Chloroplast spacer <i>trnG-trnS</i> identified five haplotype varieties under cultivation that were not present and probably were lost to extinction in the wild (Miller and Schaal et al. 2005); 216 individuals of 34 populations with two primer pair RFLPs (Miller and Schaal et al. 2006)
<i>Dimocarpus longan</i> Lour.	28 RAPD markers used to uniquely identify each of 22 accessions (163); partial <i>rbcL</i> 66 AFLP markers used on 41 <i>D. longan</i> accessions (82)
<i>Annona cherimolia</i> Mill., <i>A. muricata</i> L., <i>A. squamosa</i> L.	Development of monomorphic SSRs as phylogenetic markers of <i>Annona</i> species (Chatrou et al. 2009); 13 isozyme loci used to study segregation in 14 self-fertilized cultivars (Perfectti and Pascual et al. 1996); 206 cultivars analyzed with 23 isozyme loci in a germplasm bank (Perfectti and Pascual et al. 2005); 94 new SSRs, 58 of them polymorphic, were tested on 23 cherimoya cultivars (Escribano et al. 2008); Nine accessions discriminated by 14 polymorphic RAPD fragments (Brown et al. 2003); 5 <i>Annona</i> species analyzed with 11 allozymes indicated <i>A. muricata</i> more genetically distant from the rest (Samuel et al. 1991)
<i>Inga edulis</i>	ITS sequencing and phylogenetic study of 32 species of the seven sections of <i>Inga</i> (Richardson et al. 2001); five SSR loci to study 189 trees for possible genetic erosion (Hollingsworth et al. 2005); chloroplast <i>trnL-F</i> region and five SSR loci used to study five locations in the Peruvian Amazon (Dawson et al. 2008)
<i>Litchi chinensis</i> Sonn.	12 SSRs tested on 21 <i>Litchi</i> cultivars (159); 16 SSR markers tested on 58 <i>Litchi</i> cultivars (81)
<i>Psidium</i> spp.	23 SSRs tested on 16 accessions of <i>Psidium guajava</i> and three other species (Risterucci et al. 2005); RAPD markers associated to quercetin (Feria-romero et al. 2009); AFLP on 48 guava cultivars (Hernández-Delgado et al. 2007); revision of molecular tools applied to <i>Psidium</i> (Rai et al. 2010)
<i>Durio</i> spp. Adans.	Ten <i>Durio</i> species were analyzed for phylogenetic relationships using RFLP on two chloroplast genes <i>ndhC-trnV</i> and <i>rbcL</i> (Santoso et al. 2005); phylogenetic relationships inferred from analysis of <i>ndhF</i> and ITS sequences (Nyffeler & Baum 2000)

(continued)

Table 7.2 (continued)

Species	Work done on molecular markers
<i>Garcinia mangostana</i> , <i>G. cochinchinensis</i> , <i>G. portoricensis</i> , <i>G. binucao</i>	Random amplified DNA fingerprinting (RAF) used on 37 accessions of <i>G. mangostana</i> (Ramage et al. 2004); 21 trees analyzed with 5 primers for RAPD analysis (Sobir et al. 2007)
<i>Morinda citrifolia</i>	11 individuals of 3 varieties of <i>M. citrifolia</i> analyzed to determine geographical origin by using nrETS, nrITS, rps16 and trnT-F sequence data (Razafimandimbison et al. 2010)
<i>Adansonia</i> spp. L.	AFLP analysis of 137 individuals from six populations (Assogbadjo et al. 2006); 18 SSR markers to analyze 214 individuals of <i>Adansonia digitata</i> and 30 individuals of other <i>Adansonia</i> species (Larsen et al. 2009); 11 populations of 4 countries in West Africa analyzed by AFLP (Kyndt et al. 2009); phylogeography of 344 individuals from 74 populations analyzed by chloroplast DNA (Tsy et al. 2009)
<i>Manilkara zapota</i> (L.) van Royen	Analysis of four populations of <i>M. zapota</i> from Mexico using 4 polymorphic RAPD markers (Heaton et al. 1999); 12 polymorphic SSR loci developed for <i>M. huberi</i> (Azevedo et al. 2005)
<i>Averrhoa carambola</i> L.	None found
<i>Anacardium occidentale</i>	Fingerprinting of 19 <i>A. occidentale</i> accessions using 50 RAPD primers, 12 ISSRs, and 6 AFLPs (Archak et al. 2003); 91 individuals of <i>A. occidentale</i> analyzed by AFLP (Archak et al. 2009); 21 polymorphic SSRs from <i>A. occidentale</i> tested on other <i>Anacardium</i> species (Croxford et al. 2006)
<i>Tamarindus indica</i> L.	RAPDs used to analyze ten populations of <i>T. indica</i> (Diallo et al. 2007)
<i>Irvingia</i> spp.	Eight CAPS primers used to analyze <i>I. gabonensis</i> and <i>I. wombolu</i> (Lowe et al. 1998); 130 individuals of <i>I. gabonensis</i> and <i>I. wombolu</i> from West Africa were analyzed by RAPDs (Lowe et al. 2000); AFLP analysis of 15 accessions of <i>I. gabonensis</i> (Ude et al. 2006)
<i>Nephelium lappaceum</i> and <i>N. ramboutan-ake</i>	Seven SSR markers from <i>Litchi chinensis</i> had amplification in <i>N. ramboutan-ake</i> (Sim et al. 2005)
<i>Pometia pinnata</i>	None found
<i>Artocarpus altilis</i> (Parkinson), <i>A. heterophyllus</i> Lam.	26 <i>A. heterophyllus</i> accessions analyzed by AFLP using 12 primer pairs (136); Six populations of <i>A. altilis</i> analyzed by AFLP and 15 morphological traits (Sreekumar et al. 2007); Eight primer pairs used for AFLP analysis of 50 <i>A. heterophyllus</i> accessions (Shyamamma et al. 2008); AFLP analysis of 20 breadfruit samples of <i>A. camansi</i> and <i>A. mariannensis</i> (166)
<i>Melicoccus bijugatus</i> Jacq.	None found
<i>Syzygium samarangense</i>	Isozymes used on a related rainforest species <i>S. nervosum</i> (Shapcott et al. 1999); 8 SSRs developed for <i>S. sayeri</i> (Hillyer et al. 2007)
<i>Premna serratifolia</i> L.	None found
<i>Sandoricum koetjape</i>	None found
<i>Pouteria sapota</i>	20 cultivars of <i>M. zapota</i> analyzed by RAPD markers (Meghala et al. 2005)
<i>Dracontomelon vitiense</i>	None found
<i>Crataeva speciosa</i> Volkens	None found

The order of species is the same as in Table 7.1, in which NCBI entries are summarized

modification to the DNA extraction method, given the presence of copious latex and phenolics in the vegetative tissues. Second, for generating SSR-enriched libraries, the method previously developed by Tehen et al. (2010) was modified to adapt to high-throughput pyrosequencing with a Roche 454 GS-FLX (F. Hoffmann-La Roche Ltd., Basel, Switzerland). One modification employs two adapters (Tehen et al. 2010) that allow simultaneous loading of pairs of samples in the same region of picotiter plates; the adapters act as bar coding to separate the samples via bioinformatics. Another modification is reducing the number of PCR cycles during library preparation to minimize redundant sequences. SSRs were isolated from the following species/crop groups: *Nephelium lappaceum* Linn. (rambutan), *Manilkara zapota* Linn. (sapodilla), *Pouteria sapota* Jacq. (sapote), *Litchi chinensis* (lychee), *Melicoccus bijugatus* Jacq. (Spanish lime), *Annona squamosa* Linn. (sugar apple), *Dimocarpus longan* (longan), *Averrhoa carambola* Linn. (star fruit), *Artocarpus altilis* (Parkinson) Fosberg (breadfruit), and *Garcinia mangostana* (mangosteen).

For the combined species, a total of 2,510,291 reads were assembled into 224,815 contigs in which a total of 49,898 SSR repeats (not including mononucleotides) were detected. A total of 10,310 primer pairs were designed under stringent conditions (Tm 65°C and 3' GC clamp). Only 384 primer sets were tested per species. This was done to identify and select the SSR markers that were most readily amplifiable, reproducible, and which detected the highest allelic diversity for variety/cultivar identification and plant breeding. The SSR development process was accomplished in a short amount of time (less than 6 months), and greater than 96% of the markers resulted in SSR amplification. A detailed protocol for the SSR development procedure is available in Methods in Molecular Biology, Humana Press (Arias et al. 2010).

Specific Examples of Genomics in Tree Fruits

Avocado

The avocado (*Persea americana*) is an evergreen subtropical tree that is native from Mexico to northern South America and produces a fruit that is unique and nutritious. This fruit was known by the Aztecs as “ahuacacuahitl,” which was later shortened by the Spaniards to “aguacate.” In the United States, avocado was introduced to Florida in 1833, California in 1848, and Hawaii by 1855 (Nakasone and Paull 1998). Major commercial production of avocado in the United States is limited to California and Florida. In 2000, global production exceeded 2.4 MMT, and the major producers were Mexico, Indonesia, South Africa, and the USA (Anonymous 2002).

Persea americana has been subdivided into three horticultural groups: Mexican (*P. americana* var. *drymifolia* (Schecht. & Cham.) Blake), Guatemalan (*P. americana* var. *guatemalensis* Wms.), and West Indian (*P. americana* var. *americana* Mill.).

racess. The West Indian race is known to be from the lowland areas of the Pacific coast of Central America and not the West Indies, while the Guatemalan and Mexican races are native to specific highland areas within each country (Scora and Bergh 1992). The three racial groups can be distinguished by the percentage of oil content in the fruit, with the West Indian cultivars ranging from 2.5% to 8.0%, Guatemalan accessions from 10% to 13%, and Mexican accessions ranging from 15% to 20% (Knight 2002). The racial classes also vary phenotypically for characters such as fruit size and shape, skin thickness, skin color, seed size, and fruit ripening (Lahav and Lavi 2002). Sterility barriers do not exist between or among the three racial types (Lahav and Lavi 2002). Avocado has a distinct flowering habit known as protogynous, diurnally synchronous dichogamy (Bergh 1969). This type of reproductive behavior promotes outcrossing; however, significant amounts of self-pollination are known to occur in commercial plantings (Davenport et al. 1994; Borrone et al. 2008; Schnell et al. 2009). Named cultivars often originate from open-pollinated seedlings. The unknown pollen parent has often been estimated based on the flower types of available donor trees. Morphological characters have been used to infer parentage, although these are influenced by environmental factors and may not unambiguously distinguish closely related genotypes or interracial hybrids. Many of the cultivars grown in Florida are interracial hybrids between Guatemalan and West Indian types, while those grown in California are hybrids between Mexican and Guatemalan types.

The haploid genome size of avocado has been estimated to be 8.83×10^8 bp (Arumuganathan and Earle 1991a). Avocado contains 24 chromosomes with bivalent pairing at meiosis indicating that $n=12$ (Darlington and Wylie 1945). Avocado has been proposed to be the result of an ancient polyploid event (Chanderbali et al. 2008). However, evidence from analyses of germplasm collections and seedling populations using SSR markers has demonstrated diploid inheritance for most markers, although some primer pairs have amplified more than one locus (Schnell et al. 2003; Borrone et al. 2007, 2008, 2009).

Gene Discovery

Recently, a large number of sequences have been generated from the transcriptome of *P. americana* from two projects, the Floral Genome Project (Albert et al. 2005) and the Ancestral Angiosperm Genome Project (Wall et al. 2008). The Floral Genome Project, initiated to investigate the evolutionary development of floral patterning (Chanderbali et al. 2009), produced two directionally cloned, non-normalized, Sanger-sequenced cDNA libraries and are deposited in the EST database of GenBank (16,558 sequences). Both libraries, Pam01 (8,735 sequences) and Pam01b (7,823 sequences), were developed from premeiotic flower buds. The Pam01 library was the source of the EST-SSRs used to evaluate outcrossing rates (Borrone et al.

2007, 2008; Schnell et al. 2009) and to develop a moderately dense genetic map of avocado (Borrone et al. 2009).

The Ancestral Angiosperm Genome Project continued where the Floral Genome Project left off using pyrosequencing (Wall et al. 2008). A total of 12 libraries, developed from various tissue types and stages, generated 1,698,670 “passing” sequences covering 490.7 Mb. The unprocessed, unfiltered, raw data is deposited in GenBank as 14 Sequence Read Archives (SRAs). The processed (filtered, trimmed, and quality checked) sequence information is readily obtained from the Ancestral Angiosperm Genome Project website (<http://ancangio.uga.edu/content/aagp-home>). The website includes a detailed description of each library, a number of tools allowing comparison of the libraries with one another, and retrieval of individual reads and contigs. The entire dataset derived from pyrosequencing has been assembled with the cDNA libraries (Pam01 and Pam01b, developed from the Floral Genome Project) in a “Sanger-454 hybrid” assembly of 115,151 contigs. Polymorphic regions, SNPs and SSRs, have been identified in the assembled sequences as part of the pipeline process. The entire assembly is available for download at the site. Additional cDNA libraries have been developed from *Persea americana* var. *drymifolia* but are not yet publically available (López-Gómez et al. 2007).

An additional resource, not yet deposited in the NCBI database, is sequences of small RNAs (Blake Myers and Pamela Green, Delaware Biotech Institute, University of Delaware; NSF Plant Genome Program Award #0638525). Small RNAs, 21–24 nucleotides long, consist of two major types known as small interfering RNA (siRNA) and microRNA (miRNA) and are implicated in regulation of gene transcription/translation. These sequences were generated by pyrosequencing (Accerbi et al. 2010). A total of 12,689,456 sequences, 5,918,358 identified as being distinct, have been generated from three separate libraries: leaves (3,800,961 total; 1,263,610 distinct), flower (5,325,501 total; 2,870,885 distinct), and fruit (3,562,994 total; 1,783,863 distinct). The sequences are available through the Comparative Sequencing of Plant Small RNAs website (<http://smallrna.udel.edu/>).

Marker Development

Previously, there was a limited set of SSR markers for *P. americana* (Sharon et al. 1997; Ashworth et al. 2004), some of which do not consistently amplify in all varieties (Ashworth and Clegg 2003; Ashworth et al. 2004). For example, 14 of 39 SSR markers tested were suitable to fingerprint diverse collections of *P. americana* (Schnell et al. 2003). To increase the number of informative SSR markers, publicly available *P. americana* ESTs were screened (Borrone et al. 2007, 2009) bringing the number of markers available to upwards of 300. Data-mining SSRs from expressed sequence tags (ESTs) has proven effective for generating markers for fingerprinting, genetic mapping, and comparative mapping among species (Varshney et al. 2005).

Linkage Mapping

The first linkage map of avocado was produced by Sharon et al. (1997) from 50 progeny of a cross between 'Pinkerton' and 'Ettinger' using 50 SSR markers, 17 random amplified polymorphic DNA (RAPD) markers, and 23 minisatellite DNA fingerprint (DFP) markers. Twelve linkage groups with 34 mapped loci covering 352.6 centimorgans (cM) were identified. Seven linkage groups contained two markers, two linkage groups contained three markers, one linkage group contained four markers, and two linkage groups contained five markers each. A larger population and an increased number of genetic markers were needed to produce a linkage map useful for quantitative trait loci (QTL) discovery. The development of over 300 SSR markers (Sharon et al. 1997; Ashworth et al. 2004; Borrone et al. 2007, 2009) enabled the development of a partially saturated genetic linkage map and the potential identification of QTLs controlling horticultural traits of interest in avocado. Mapping populations have been developed in California (Ashworth et al. 2007; Chen et al. 2007) and in Spain (Viruel et al. 2007) by producing full-sib families that are similar in size to those used by Sharon et al. (1997) and are focused upon Guatemalan-Mexican hybrids.

In Florida, a large population of seedlings from a commercial grove interplanted with two cultivars of opposite flowering types, 'Tonnage' (type B) and 'Simmonds' (type A), in approximately equal numbers was screened to determine the outcrossing rate in avocado under south Florida conditions (Borrone et al. 2008). Eight fully informative SSR markers identified 870 seedlings as progeny of a reciprocal cross between 'Tonnage' and 'Simmonds'. Using these seedlings, the first large mapping population and linkage map for QTL discovery were developed in avocado, focusing upon West Indian-Guatemalan hybrids (Borrone et al. 2009). The final linkage map (Fig. 7.1, Table 7.3) was constructed from 163 markers generated by 135 primer pairs, 112 designed from EST-SSRs and 23 SSR primers previously developed by Sharon et al. (1997). Twenty primer pairs amplified more than one locus, with 15 amplifying two loci, four amplifying three loci, and one amplifying six loci. 'Tonnage' was heterozygous for 92.0% of the total markers used in its sub-composite map, whereas 'Simmonds' was heterozygous for 41.0% of the total markers in its sub-composite map, comparable with the previous estimate for heterozygosity for each cultivar using 12 SSR markers (Borrone et al. 2008). Eighteen markers were fully informative with four polymorphic alleles between the two parents, and the other 145 markers were partially informative. One hundred nineteen markers were scored on both sub-composite maps. An additional 43 markers were scored only on the 'Tonnage' × 'Simmonds' progeny, and one additional marker was scored only on the 'Simmonds' × 'Tonnage' progeny.

Twelve linkage groups (LG) representing the haploid set of 12 chromosomes of *P. americana* were generated from 163 markers, at a minimum logarithm (base 10) of odds (LOD) score of 4.0 (Fig. 7.1, Table 7.3). Linkage groups ranged in size, longest to shortest, from 157.3 (LG2) to 2.4 cM (LG12), and the number of markers mapped per group ranged from 29 (LG1) to 2 (LG12). The total length of the Florida

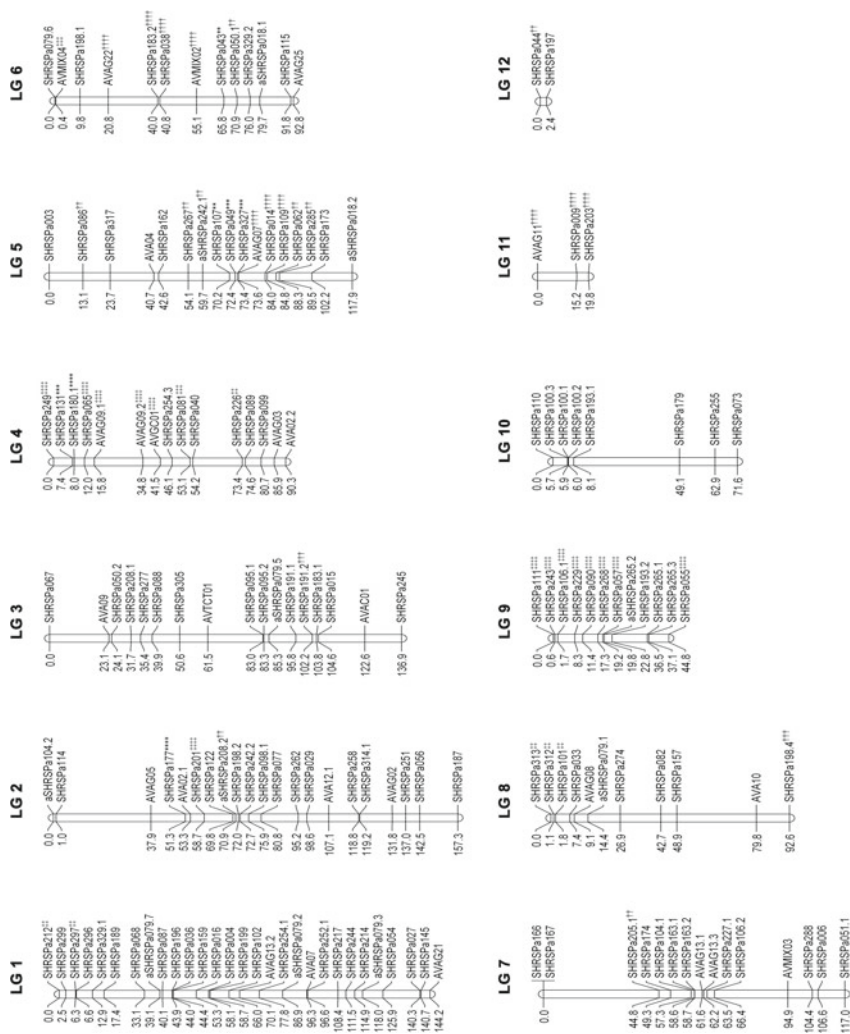


Fig. 7.1 Avocado linkage map constructed from 163 markers generated by 135 primer pairs

Table 7.3 *Persea americana* linkage mapping results for the Florida mapping population. Mapping results of the reciprocal cross (sub-composite maps) and parental maps

Map	LG	1	2	3	4	5	5a ^a	5b	5c	6	7	8	9	10	11	12	Total
'Tonnage' × 'Simmonds' 456 individuals	No. loci	29	21	16	15	12	3	2	13	15	11	11	12	8	3	2	162
	Length (cM)	143.9	170.1	138.7	95.7	117.3	17.6	29.3	93.7	119.6	95.3	46.6	71.3	23.7	3.1	1,119.0	
	cM/locus	5.0	8.1	8.7	6.4	9.8	5.9	14.7	7.2	8.0	8.7	3.9	8.9	7.9	1.6	7.5	
'Tonnage' maternal	No. loci	27	18	16	15	15			12	13	11	11	8	3	2	149	
	Length (cM)	146.5	154.7	138.6	96.3	116.5			92.1	121.1	95.4	47.0	70.9	23.8	3.1	1,106	
	cM/locus	5.4	8.6	8.7	6.4	7.8			7.7	9.3	8.7	4.3	8.9	7.9	1.6	7.1	
'Simmonds' paternal	No. loci	16	11	3	4	3	3		7	5	0	6	2	0	0	60	
	Length (cM)	144.8	113.6	38.5	35	19.9	41.9		100.8	68.6	–	48.4	46.9	–	–	658.4	
	cM/locus	9.1	10.3	12.8	8.8	6.6	14.0		14.4	13.7	–	8.1	23.5	–	–	12.1	
'Simmonds' × 'Tonnage' 259 individuals	No. loci	20	16	11	13	10	2		9	12	8	8	6	3	2	120	
	Length (cM)	141.1	132.2	115.5	79.6	84.4	14.8		89.2	99.1	74.6	39.8	68.9	13.0	1.2	953.4	
	cM/locus	7.1	8.3	10.5	6.1	8.4	7.4		9.9	8.3	9.3	5	11.5	4.3	0.6	7.4	
'Simmonds' maternal	1a																
	1b																
	No. loci	11	2	8	0	4	6		6	3	0	5	0	0	0	0	45
'Tonnage' paternal	Length (cM)	69.1	16.5	92.6	–	41.3	103.6		92.3	42.6	–	44.3	–	–	–	–	502.3
	cM/locus	6.3	8.3	11.6	–	10.3	17.3		15.4	14.2	–	8.9	–	–	–	–	11.5
	No. loci	18	14	11	13	10			9	11	8	8	6	3	2	113	
'Tonnage' maternal	Length (cM)	141.1	124.6	115.4	77.5	106.3			85.6	99.5	74.6	37.1	68.8	13	1.2	944.7	
	cM/locus	7.8	8.9	10.5	6.0	10.6			9.5	9.0	9.3	4.6	11.5	4.3	0.6	7.7	

^aLG 5a represents the entire length of the LG in the 'Tonnage' × 'Simmonds' map and was used to calculate the overall length of the linkage map. For all others, LG5 are smaller-sized fragments of the entire LG

F_1 map was 1,087.4 cM. This Florida F_1 map is three times the size of the map reported by Sharon et al. (1997) and contains five times as many markers. Comparison of markers shared between the Florida F_1 map and the F_1 map of Sharon et al. (1997) allowed the identification of several analogous linkage groups. Ten linkage groups of the 12 described by Sharon et al. (1997) contained SSR markers. Nine of the ten corresponded to seven linkage groups in the Florida F_1 map. Distances between SSR markers within LGs reported by Sharon et al. (1997) and distances between these same markers placed in the sub-composite maps and the Florida F_1 map corresponded well, which is remarkable given the disparity in the population sizes and the numbers of markers between the two maps. The ‘Tonnage’ \times ‘Simmonds’ linkage groups were longer than the ‘Simmonds’ \times ‘Tonnage’ linkage groups (Table 7.3). This can be attributed to the larger number of markers mapped in the ‘Tonnage’ \times ‘Simmonds’ sub-composite map versus its reciprocal (162 vs. 120) and to the larger number of individuals in this cross (456), slightly less than double that of the ‘Simmonds’ \times ‘Tonnage’ cross (259). Also, LG5 formed three linkage groups in the ‘Tonnage’ \times ‘Simmonds’ map and two linkage groups in the ‘Simmonds’ \times ‘Tonnage’ map. The parts of this linkage group reformed a single linkage group in the composite map. Five linkage groups (LGs 3, 4, 8, 11, and 12) were composed almost entirely from markers polymorphic in the ‘Tonnage’ parent. The formation of linkage groups for all maps was straightforward except for a small number of markers due to the high homozygosity of ‘Simmonds’.

The number of SSR markers now available for avocado allowed the development of this first moderate-density map composed of 163 loci. Linkage maps developed using molecular markers, such as this Florida F_1 map, enable the detection and use of QTLs affecting traits of economic importance. The two parental cultivars ‘Simmonds’ and ‘Tonnage’ differ for many phenotypic traits useful in humid low-land environments like south Florida. ‘Simmonds’ is of the West Indian race and believed to be a seedling of ‘Pollock’. It was selected in south Florida, propagated commercially in 1921, and is still a major commercial cultivar some 90 years later. ‘Simmonds’ has a light green, oblong-oval to pear-shaped large fruit with medium-sized seed and low oil content (3–6%), imparts tolerance to *Phytophthora* root rot (PRR) in its progeny, has the “A” flowering type, and is a high-yielding, early-season (June–July) cultivar (Campbell and Malo 1978; Ploetz et al. 2002). West Indian selections were the only important commercial cultivars in Florida until the 1920s when competition from Cuba depressed the market for Florida avocados. A number of Guatemalan–West Indian hybrids had since been selected which ripened in the fall and winter, extending the season and plantings shifted to include these hybrids. One of these was the cultivar ‘Tonnage’ which is now considered a minor cultivar in south Florida. ‘Tonnage’, a seedling of ‘Taylor’, was first propagated commercially in 1930. It has a dark green, pear-shaped fruit with medium seed size and moderate oil content (8–15%), does not impart tolerance to PRR in its progeny, is a “B” flowering type, and is a late-season (August–September) cultivar (Ploetz et al. 2002).

Recently, the USDA-ARS Subtropical Horticulture Research Station (SHRS) in Miami, FL, has produced another large mapping population of ‘Hass’ \times ‘Bacon’ (~1,000 individual seedlings). ‘Hass’ is the most important avocado cultivar worldwide (Ashworth et al. 2007), and growers receive a premium price for ‘Hass’ fruit.

A commercial California ‘Hass’ avocado orchard was identified with adjacent ‘Bacon’ pollinizer rows. ‘Hass’ is an interracial hybrid of mostly Guatemalan ancestry, and ‘Bacon’ is of the Mexican landrace; they are of opposite flowering types (A and B, respectively) and, thus, expected to outcross. The full-sib family members from this ‘Hass’ × ‘Bacon’ cross were identified by genotyping with five SSR markers, and seedling trees, product of self-pollination, were excluded. However, a map has not yet been produced. Presently, the limiting factor is the availability of sufficient molecular markers to make a saturated, high-resolution map for QTL mapping of important agronomic traits such as oil content and composition, flowering type, flowering time, and cold tolerance. Once the field data for agronomic traits is collected on this population, this map will be useful for detection and use of QTLs.

SNP Discovery

SNP discovery from transcriptome and genomic sequence data is needed for obtaining sufficient molecular markers for complete linkage map saturation. The following strategy has been used at SHRS to generate and validate new SNP and SSR markers. The strategy takes advantage of two next-generation sequencing platforms, Illumina GAI (Illumina, Inc., San Diego, CA) and Roche 454, and uses both transcriptome and genomic sequence data to identify and validate SSRs and SNPs. Once sufficient SNPs have been identified (5–10 K), the production of an Illumina Infinium oligonucleotide array for genotyping of mapping populations may be employed. ‘Hass’, the most important commercially grown avocado cultivar in the world, has been selected as the reference cultivar for the transcriptome sequencing.

Transcriptome Sequencing

RNA was isolated from leaves, unopened flowers, female flowers, and male flowers from ‘Hass’, ‘Bacon’, ‘Simmonds’, and ‘Tonnage’, from tissue collected from genotyped clones at the SHRS. The RNA from the ‘Hass’ tissue was pooled for Roche 454 sequencing to generate the reference transcriptome. In addition, a large EST library was produced by Roche 454 sequencing of developing mesocarp tissue of ‘Hass’ and that data was used to expand the ‘Hass’ reference transcriptome (John Ohlrogge, Michigan State Univ., personal communication 2010). The RNA from ‘Bacon’, ‘Simmonds’, and ‘Tonnage’ leaves and flowers was pooled for each clone and sequenced by Illumina GAI. Illumina reads were aligned to the ‘Hass’ reference transcriptome, and a variant report with single-nucleotide polymorphisms and indels generated. The variant report was filtered for heterozygous SNP loci, as a locus that is heterozygous in any parent of the mapping populations can be mapped. SSR motifs are identified from the ‘Hass’ transcriptome sequences as described

earlier in this chapter. The SNP and SSR discovery will yield a sufficient number of markers to completely saturate the genetic maps for both populations.

Reduced Representational Sequencing of the ‘Hass’ Genome

Reduced representational sequencing of the genome of ‘Hass’ from hypomethylated DNA is being carried out using Roche 454 pyrosequencing to obtain a ~2–4X coverage of the avocado gene space. Hypomethylated DNA is actively transcribed, and this should enrich for gene-containing regions of the genome. In the cacao SNP and SSR discovery project (<http://www.cacaogenomedb.org>), the genomic sequence has proved essential for identifying intron-exon junctions in the transcriptome sequences prior to SNP oligonucleotide design or SSR primer design. In addition, the assembled gene space of ‘Hass’ will be made available to the avocado genome sequencing group, producing a hypomethylated library of the Mexican landrace cultivar by an alternative method. Such a comparison will be of great interest for future genome sequencing projects targeting only the gene space.

Genome Sequencing Project

The avocado genome sequencing project has been under way for over a year in Mexico at the Plant Biotechnology Unit, Centre for Research and Advanced Studies (CINVESTAV; Luis Herrera-Estrella and Gustavo Hernandez), as well as by the bio-fuels research on non-seed oils group in the Plant Lipid Metabolism lab at Michigan State University (John Ohlrogge). The Mexican avocado genome sequencing project is using a highly homozygous individual from the Mexican landrace (*P. americana* var. *drymifolia*). Using flanking sequences of the 163 SSR markers mapped in the Florida F1 population, CINVESTAV has designed overgo probes to anchor their physical map to the genetic recombination map. The current version of the avocado physical map has 5,008 contigs, and the estimated coverage is 5X. Using the 163 markers will certainly help in the anchoring process, but thousands of markers would allow anchoring and correct ordering of the entire physical map of avocado. The Illumina Infinium chip being designed from the SNP development project will saturate the avocado genetic recombination map and anchor physical map contigs.

Mango

Mango (*Mangifera indica*) is a significant tree fruit crop grown commercially in tropical and subtropical areas of many countries. It has been under cultivation in India for at least 4,000 years, and over 1,000 varieties are known to exist in that country

(Mukherjee 1953). India is the largest producer of mangos with 10.8 million MT accounting for 41% of the world's mango production (Gunjate 2009, Proc VIII int. mango sym.). Most commercial mango cultivation is in tropical and subtropical regions; however, mangos are also cultivated in a wide range of marginal climatic areas throughout the world. Mango genotypes are divided into two distinct categories or types based on their origin: monoembryonic mangos, which are mostly subtropical (Indian types), and polyembryonic mangos, which are mostly tropical (Southeast Asian types). The seeds of Indian types characteristically contain a zygote embryo, and the fruit skin is highly colored (mixes of red, purple, and yellow), while the seeds of Indo-Chinese types contain several nucellar embryos, and the skin is soft or pale in color (green to light green to yellow) (Iyer and Degani 1997; Viruel et al. 2005).

Information on the cytology of mango is quite limited. Only *Mangifera* species *M. indica*, *M. caloneura*, *M. sylvatica*, *M. foetida*, *M. caesia*, *M. odorata*, and *M. zeylanica* have been studied, and these were found to have chromosome numbers of $2n=40$ and $n=20$ (Mukherjee 1950, 1957, 1963). Chromosome numbers and ploidy status of other *Mangifera* species have yet to be studied (Bompard and Schnell 1997). Mango has a small haploid genome size (0.91 pg), which is three times larger than *Arabidopsis thaliana* (L.) Heynh. and comparable to that of rice (Arumuganathan and Earle 1991b). Mango has been referred to as an allopolyploid based on the conclusions drawn by Mukherjee (1950). Due to the presence of secondary associations at metaphase of meiosis, he suggested that the basic chromosome number of *Mangifera* is $n=8$. In addition, the high number of somatic chromosomes and the correspondingly high number of nucleolar chromosomes led him to conclude that mango is an allopolyploid. However, the evidence used to arrive at this conclusion is not unequivocal. In fact, the molecular marker evidence is antithetical to this conclusion. Results from Duval et al. (2005), Schnell et al. (2005, 2006), and Viruel et al. (2005) all resulted in normal diploid segregation for SSR markers.

Applications of Molecular Genetics

Isozymes were the first markers to be used for fingerprinting mango cultivars to determine self- vs. cross-pollination and to estimate genetic relationships (Degani et al. 1990; Knight and Schnell 1994). RAPD markers were also used to fingerprint cultivars and estimate genetic relationships in mango (Schnell et al. 1995). In that research, a group of 'Haden' seedlings and a random group of seedlings were evaluated using 11 RAPD primers. This study supported the 'Haden' parentage of 'Eldon', 'Lippens', 'Tommy-Atkins', and 'Zill'; however, the parentage of 'Glenn' and 'Osteen' was questioned. Adato et al. (1995) used DNA fingerprinting (DFP) to evaluate genetic relationships between 26 mango cultivars and 14 rootstocks. They provided a pedigree that further confirmed the relationship between many of the 'Haden' seedlings. Lopez-Valenzuela et al. (1997) used RAPD markers to estimate genetic diversity among 15 rootstock cultivars using 13 markers and identified a specific RAPD band associated only with the polyembryonic types. Eiadthong et al.

(1999) utilized anchored simple sequence repeat markers to analyze 22 mango cultivars; they were able to distinguish genotypes. However, the authors were unable to find markers unique to either monoembryonic or polyembryonic types or for the Thai cultivars selected for green harvest (crispy mango) from the cultivars selected for ripe fruit production. Pandit et al. (2007) also used inter-simple sequence repeat (ISSR) markers to evaluate 60 elite Indian mango cultivars and 10 non-Indian cultivars. They were not able to distinguish Indian cultivars from northern and southern India, and they concluded that ISSR markers could not be considered a comprehensive marker system for mango. Kashkush et al. (2001) utilized amplified fragment length polymorphisms (AFLP) to estimate genetic relationships between 16 cultivars and 7 rootstock cultivars. They also analyzed 29 progeny from a cross of ‘Tommy-Atkins’ and ‘Keitt’ and produced a crude linkage map that identified 13 of the 20 linkage groups. The start codon targeted (SCoT) markers have also proved useful in evaluating the relationship among the Xiang Ya Mango type cultivars in China (Luo et al. 2011). SCoT markers are useful for the confirmation of identical genotypes and clones with different names. SCoT and other dominant markers are useful for germplasm collection management, and they are technically simple and inexpensive to implement.

SSR Markers

Viruel et al. (2005) developed the first reported set of 16 SSR markers for mango, of which 14 produced the expected one or two amplification products per genotype. These 14 SSRs were used to evaluate 28 mango genotypes that included 14 Florida cultivars. Discrimination of all 28 genotypes was possible, and the average number of alleles per locus was 5.3. Previously known pedigree information for the ‘Haden’ family of mangos was confirmed and was in agreement with published RAPD and DFP analyses (Adato et al. 1995; Schnell et al. 1995) with one exception. Viruel’s clone of ‘Zill’ was not resolved as a seedling of ‘Haden’. Schnell et al. (2005) developed a second set of 15 SSR markers and analyzed 59 Florida cultivars and four related species. Two of the SSRs were monomorphic among the Florida cultivars; the other 13 had an average number of alleles per locus of 4.2 with polymorphism information content (PIC) values varying from 0.21 to 0.63.

Schnell et al. (2006) used 25 SSR loci to estimate genetic diversity among 203 unique mangos (*M. indica*), two *M. griffithii* Hook. f. & Thomson, and three *M. odorata* Griff. accessions maintained at the National Germplasm Repository (NGR) and at the Fairchild Tropical Garden (FTG) in Miami, Florida. The 25 SSR loci had an average of 6.96 alleles per locus and an average PIC value of 0.552 for the *M. indica* population. The total propagation error in the collection (i.e., plants that had been incorrectly labeled or grafted) was estimated to be 6.13%. When compared by origin, the Florida cultivars were more closely related to Indian than to Southeast Asian cultivars. Unbiased gene diversity (H_{nb}) of 0.600 and 0.582 was found for Indian and Southeast Asian cultivars, respectively, and both were higher than H_{nb} among Florida

cultivars (0.538). When compared by horticultural type, H_{nb} was higher among the polyembryonic types (0.596) than in the monoembryonic types (0.571).

Until recently, a total of 62 SSR markers had been developed for mango (Duval et al. 2005; Honsho et al. 2005; Schnell et al. 2005; Viruel et al. 2005). This number is more than adequate for genetic diversity studies and for parentage analysis as has been demonstrated by Schnell et al. (2006); however, these numbers are not sufficient to develop a saturated linkage map for the 20 linkage groups of mango. Over 300 new SSR markers have been developed at the USDA-ARS Mid South Area Genomics Laboratory (MSAGL) using pyrosequencing and verified using a mango diversity panel of 11 cultivars, using the method described earlier in this chapter. Another additional 1,800 SSR markers have been produced in the Australian mango genomics program (Ian Bally, personal communication 2010).

Parentage Analysis

The SSR markers that have been developed for mango are easily used to verify parentage using a software package like CERVUS (Marshall et al. 1998). When caging trees to exclude foreign pollen or using the polycross mating design, it is now simple to identify the male parent from a set of potential male parents. This has been very useful in cacao breeding where mistakes in pollination have led to the estimation of unreliable breeding values for parental clones.

Among the 64 Florida cultivars evaluated in the parentage analysis by Schnell et al. (2006), the genetic background was found to be based on as few as four Indian cultivars and the polyembryonic cultivar 'Turpentine'. Two Indian cultivars, 'Mulgoba' and 'Sandersha', are in the background of most Florida types with 'Amini', 'Bombay', 'Cambodiana', 'Long', 'Julie', 'Turpentine', and 'Nam doc Mai' making lesser contributions. The seedling races of Cuba and Florida were considered the same by Popenoe (1920) who called them the West Indian race commonly known as 'Turpentine' in Florida. In the parentage analysis, 'Turpentine 10' was identified as a most probable paternal parent for 'Haden'. 'Haden' was reported as the maternal parent for 10 cultivars included in the analysis, but based on the parentage analysis, 31 cultivars were found to have 'Haden' as one of the most likely parents. Likewise, the other important early Florida selection 'Brooks' is the parent of seven cultivars. 'Haden', 'Brooks', and seedlings of 'Haden' and 'Brooks' have contributed disproportionately to the genetic background for many of the cultivars in the Florida group.

Linkage Mapping

The first genetic linkage map in mango was reported by Kashkush et al. (2001), utilizing AFLP markers and 29 progeny from a cross of 'Tommy-Atkins' × 'Keitt' in Israel. They were able to map 34 AFLP loci and produced a crude linkage map that

identified 13 of the 20 linkage groups covering 160 cM. A second map has been produced using 60 progeny from a cross of 'Keitt' × 'Tommy-Atkins' in China using AFLP markers. Eighty-one markers with the correct segregation ratios were identified, and 39 of these were used to identify 15 linkage groups. The average distance between two adjacent markers was 14.74 cM. Improvement of the mango recombination map requires the development of more codominant molecular markers. Using Roche 454 sequencing and the SSR discovery pipeline discussed earlier in this chapter, new SSR markers have been developed and verified and are now being validated at SHRS on mapping populations. A suitable number of SSR markers are being identified to develop a moderately saturated recombination map for mango.

Florida Mapping Populations

Two experimental populations have been developed and planted in the field at SHRS as mapping populations. The first population is an F_2 population derived from self-pollination of 'Tommy-Atkins' consisting of 168 seedlings that were planted in the field in 1995. The second population is an F_2 population derived from self-pollination of 'Haden'. A total of 224 seedlings from a single isolated 'Haden' tree have been in the field for 3 years. Phenotypic data collection is in progress for both of these populations. The development of a saturated linkage map and the identification of QTL for important traits are objectives for the USDA-ARS program at SHRS in Miami for the next 5 years.

Australian Mapping Populations

In the Australian program, supported by Agri-Science Queensland, three mapping populations have been developed using controlled pollinations (Ian Bally, Queensland DEEDI, personal communication). The largest populations are 'Irwin' × 'Kensington Pride', 'Tommy-Atkins' × 'Kensington Pride', and 'Creeper' × 'Kensington Pride'. 'Kensington Pride' is the primary commercial mango cultivar in Australia and has a fine flavor and good productivity. A consensus linkage map has been produced using 700 AFLP and 10 SSR markers. Phenotypic data is currently being collected on the mapping populations for QTL analysis, and additional SSR markers are being added to the maps. These maps have not been published.

Marker-Assisted Selection

To develop a marker-assisted selection (MAS) program for mango, more extensive linkage maps need to be developed and mapping populations field evaluated. The progeny size of both of the Florida mapping populations is small (168 and 224). In

the Australian program, sizable populations have been developed from controlled pollination. These populations are the best candidates for QTL identification in mango. The populations developed from isolated groves by the USDA-ARS SHRS in Florida together with the Australian populations will be used in the next few years to produce a comprehensive linkage map and to identify QTL for disease resistance and for important horticultural traits.

Holton (2010) reported that the Queensland Primary Industries and Fisheries has invested in a gene discovery project for mango with the goal of discovering genes controlling consumer and grower traits, such as fruit quality and tree architecture. Using a multidisciplinary approach, they are sequencing expressed genes via ESTs and serial analysis of gene expression (SAGE), using next-generation sequencing producing low-pass genome converge, identifying candidate genes from fruit quality and tree architecture, and identifying aroma volatiles from fruit.

Lychee

The lychee, *Litchi chinensis*, is the sole member of the genus *Litchi* in the soapberry family Sapindaceae. It is a tropical fruit tree native to southern China, which is also the largest lychee-producing country and the center of diversity for this genus. It is a diploid with $x=15$ and $2n=30$. A number of manuscripts have been published using RAPD and AFLP markers to estimate genetic diversity in lychee including, Tongpamanak et al. (2002) working with Thai cultivars, Kumar et al. (2006) with Indian cultivars, and Jones et al. (2006) working with the USDA-ARS germplasm collection at the Pacific Area Basin Research Center (PBARC) in Hilo, Hawaii. Substantial genetic diversity has been found among cultivars and a high level of heterozygosity detected in these studies. A limited number of SSR markers were developed for lychee (Viruel and Hormaza 2004; Li et al. 2006) and used to evaluate germplasm collections. In contrast to the RAPD and AFLP marker studies, Viruel and Hormaza (2004) analyzed 21 lychee cultivars using SSR markers, and their results suggested a very narrow genetic base. Li et al. (2006) also found limited genetic diversity when evaluating 58 lychee cultivars from Hainan Island using SSR markers. Using the SSR-enrichment method of Tehen et al. (2010) in combination with pyrosequencing, 384 markers were developed and are being verified for this species.

Linkage Map

A high-density recombination map was constructed using AFLP and RAPD molecular markers using 73 individuals from a cross of 'Maguili' \times 'Jiaohesanyuehong' (Liu et al. 2010). These individuals are highly divergent, as 'Maguili' is a very late ripening cultivar, while the male 'Jiaohesanyuehong' is a very early ripening cultivar.

Using AFLP and RAPD markers that segregated in the double pseudo-testcross configuration, 425 maternal specific markers, 357 paternal specific markers, and 257 that were segregating in both parents were identified. Using JoinMap 3.0, a map was developed with 312 markers, covering a total genetic distance of 1,040 cM and identifying 16 linkage groups. Among the linkage groups, MS1 and MS2 had a large number of markers relative to the other linkage groups, 102 and 76, respectively. The remaining 14 linkage groups contained between 12 and 4 markers (Liu et al. 2010). The progeny of this cross were also evaluated for the length of the juvenile phase and for fruit maturation date and fruit growth duration. Twenty markers closely related to fruit maturation were discovered; however, QTLs were not identified for any of the traits (Fu et al. 2008).

Longan

Longan, *Dimocarpus longan*, is native to southern China in Yunnan province and is an important tree fruit crop in the Sapindaceae family. It is a diploid with $x=15$ and $2n=30$. The fruit has a bright black seed, and the literal meaning of the word longan is “eye of the dragon.” Isozymes were first used to identify longan cultivars in the 1980s (Chen and Ke 1989); however, this method was not very useful in distinguishing cultivars. RAPDs were the first PCR-based marker developed for longan, allowing the separation of different groups and the identification of cultivars (Chen and Liu 2001; Lin et al. 2005; Zhong et al. 2007). Yonemoto et al. (2006), using RAPD markers, was able to distinguish three different groups of germplasm from Taiwan, China, and Malaysia. AFLPs were used by Yi et al. (2003), and they were able to distinguish 11 clusters among the 46 cultivars studied. ISSR markers (Hong 2007) have also been developed and used for cultivar identification and diversity analysis. Using the SSR-enrichment method of Tehen et al. (2010) in combination with pyrosequencing, 384 markers were developed and are being verified for this species.

Linkage Map

Guo et al. (2010) constructed a linkage map using 94 progeny from a cross of ‘Fengliduo’ and ‘Dawuyuan’. ‘Fengliduo’ is a high-quality cultivar, while ‘Dawuyuan’ has a large fruit size. Using RAPD, ISSR, sequence-related amplified polymorphism (SRAP), and AFLP markers in a pseudo-testcross design, JoinMap 3.0 was used to construct the linkage maps. For the cultivar ‘Fengliduo’, 183 loci were mapped and 21 linkage groups were identified covering 965 cM. For the cultivar ‘Dawuyuan’, 251 markers were mapped into 22 linkage groups covering 1,064 cM. This is the first reported map on longan which needs to be enhanced with codominant markers.

Summary

The ease of development of molecular markers, especially those amenable to high throughput in a MAS program will greatly enhance the rate of genetic gain for many of these tropical fruit species. The development of second- and third-generation pyrosequencing combined with the SSR and SNP discovery pipelines has removed many of the constraints on the use of molecular markers in these tropical fruit species. Thousands of SSR markers and hundreds of thousands of SNP markers can now be discovered with just a few Roche 454 or Illumina runs. With the removal of the limitation on numbers of markers, much of the difficulty in applying these techniques has been greatly reduced. The more challenging problem is the development of large mapping populations and the accumulation of phenotypic data to identify QTL regions associated with traits of interest. Once these populations are made and evaluated, the application of MAS to tropical fruit crops is straightforward.

Complete genome sequencing for most of these species is still a distant possibility. *De novo* assembly of second-generation sequence data into pseudo-molecules representing complete chromosomes requires a significant investment of time and funds. Considering the limited commercial value of many of these tropical species, complete genome sequencing and assembly are not likely using current technology. However, the ability to generate thousands of markers and to apply MAS will greatly accelerate tropical fruit breeding programs.

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